WO 2004/054576 PCT/EP2003/014422

PHARMACEUTICAL COMPOSITIONS OF SERTACONAZOLE FOR VAGINAL USE

Field of the invention

The present invention relates to compositions of sertaconazole for vaginal use and more specifically to compositions of sertaconazole for vaginal use in the treatment of vulvovaginal candidiasis.

Background of the invention

Vulvovaginal candidiasis is an inflammatory process that affects the vulva, the vagina or both together, and is caused by a superficial infection of the epithelial cells, especially by the yeast Candida albicans and to a lesser extent by other Candida spp., such as C. glabrata, C. tropicalis, C. parapsilosis, C. guillermondi and C. krusei. Vulvovaginal candidiasis is characterised by pruritus, vaqinal secretion with or without true vaginitis, leucorrhoea, vulvar erythema, and maceration. prevalence of this disease is increasing, the research for and development of new antifungal preparations is fully nowadays accepted justified. It is that oral intravaginal antifungal drugs are similarly effective in the treatment of uncomplicated vulvovaginal candidiasis. Since it is usually preferable to administer medicines topically rather than orally, especially in pregnant women, local treatment of vulvovaginal candidiasis is consequently recommended and oral drug delivery should be avoided whenever possible.

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USP 4551148 describes systems for vaginal delivery consisting of emulsions or suspensions of nystatin with characteristics of bioadherence to the vaginal surface. On the other hand, US Patent 5266329 describes systems for vaginal delivery consisting of emulsions or suspensions of imidazole antifungal agents with characteristics of bioadherence to the vaginal surface.

WO95/31178 describes emulsions and aqueous solutions of itraconazole with cyclodextrin for vaginal use.

USP 5514698 describes long-lasting antifungal vaginal creams that have a stable viscosity in the human body.

EP 770384 describes anhydrous solid pharmaceutical compositions of antimycotic agents, antiprotozoal agents, disinfectants, hormones, antibiotics and chemotherapeutic agents for vaginal use containing polycarbophil as a unique mucoadhesive polymer. Similarly, EP 918510 describes gels of polycarbophil-azole complexes with antifungal, or antiprotozoal activity, in which the polycarbophil acts as a mucoadhesive polymer.

WO00/30626 describes a method for treating vulvovaginal candidiasis consisting of the intravaginal administration of a single dose of an ovule of miconazole nitrate as well as the application of miconazole nitrate cream to the vulva.

30 WO02/03896 describes a method for treating vaginal or uterine infections caused by fungi, bacteria, viruses or parasites that consists of bringing the vaginal epithelium into contact with an intravaginal device that contains an

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antifungal agent, an antibacterial agent, an antiviral agent or a trichomonocidal agent, including a lipophilic or hydrophilic excipient, a mucoadhesive agent and a penetration enhancer of the active ingredient.

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DE-A-19737348 describes synergistic combinations of clindamycin and clotrimazole in the form of tablets, pessaries and ovules for local treatment of bacterial and fungal infections of the vagina.

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WO 99/55333 describes synergistic combinations of at least two imidazole ingredients for locally combating the microorganisms that cause vulvovaginitis and vaginosis.

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WO 03/032948 describes compositions and methods for delivering antibacterial, antifungal and antiviral ointments to the oral, nasal or vaginal cavity. Said compositions may contain one or more bioadhesive agents, such as xanthan gum and sodium carboxymethylcellulose.

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pharmaceutical compositions to 99/13862 relates WO pharmaceutically acceptable bioadhesive comprising carrier for vaginal administration. The bioadhesive carrier is a cross-linked polycarboxylic acid polymer formulation. Suitable cross-linking agents include divinyl divinylbenzene, N, N-dialkylacrylamide, 3,4-hydroxy-1,5hexadiene, 2,5-dimethyl-1,5-hexadiene and similar agents. Suitable polycarboxylic acids include polyacrylic and polymethacrylic acids.

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In Torres et al., International Journal of Gynecology & Obstetrics 71 (2000) 53-520 the use of sertaconazole in gynecology is described. The formulations and doses tested

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were 300 mg sustained release vaginal ovule, 500-mg vaginal tablet (both single dose) and 2 % vaginal cream in repeated applications for 7 days.

Detailed description of the invention

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The object of the present invention is to provide new compositions of sertaconazole for vaginal use in the treatment of vulvovaginal candidiasis. More specifically, the present invention relates to mucoadhesive vaginal compositions of sertaconazole in single-dose dosage forms for the treatment of vulvovaginal candidiasis.

No composition of sertaconazole with the aforesaid characteristics has been described to date.

Sertaconazole is a broad-spectrum antifungal agent with excellent activity against yeasts, dermatophytes, addition to its antifungal opportunistic In fungi. profile, safety good sertaconazole has а efficacy, sustained cutaneous retention and low systemic absorption. All these properties make it be an ideal product for For reference, the intopical application. activities, expressed as minimum inhibitory concentrations (MIC), of sertaconazole, bifonazole and econazole against mostly prevalent Candida spp. in vulvovaginal candidiasis are shown in Table 1 (Carrillo-Muñoz A.J. and Torres-Rodriguez J.M., J. Antimicrob. Chemother. 1995: 36, 713-716).

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Table 1

Microorganism	Sertaconazole	Bifonazole	Econazole
C. albicans (73)	1.02	3.6	2.24
C. tropicalis (21)	1.67	9.51	3.14
C. glabrata (16)	0.78	4.09	2.39
C. parapsilosis (22)	0.31	3.76	0.75
C. krusei (13)	0.38	2.20	0.91
C. guillermondii (5)	0.51	3.87	1.11

Furthermore, sertaconzole is superior to most imidazole antifungal drugs as a fungicide against *C. albicans* (Palacín C., Sacristán A. and Ortiz J.A., Arzneim. Forschung, 1992: 42(I), 711-714; Agut J., Palacín C. and Ortiz J.A., Arzneim. Forschung 1992, 42(I), 721-724).

compositions, the present contrast to prior-art In invention is characterised by the presence of one or more mucoadhesive excipients. These mucoadhesive excipients are preferably selected from cellulose polymers, such as methylcellulose, carboxymethylcellulose, hydroxypropyl methylcellulose and the like, or from polyacrylic acidderivative polymers, such as carbomers, polycarbophils and discovered The applicants have like. surprisingly, the combination of a polycarbophil and a the mucoadhesive action enhances carbomer preparation, but not the absorption of sertaconazole. Consequently, the active ingredient, sertaconazole, remains in the mucosa of the vagina for a period of 3 to 5 days,

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its absorption (permeation) through the vaginal mucosa being less than 0.1% of the dose. As a result, systemic side effects are negligible. The resulting intravaginal preparation requires just a single-dose application to achieve a convenient and safe eradication of *Candida* spp., which is very advantageous in practice.

The excipients used in the present invention are classified as lipophilic, mucoadhesive and preservative. Among the possible excipients, which are not intended to restrict the scope of the present invention, the following are preferred:

- stearates and excipients: Glyceryl Lipophilic a) for example, polyethylene derivatives, stearates, ketostearyl alcohols, polyoxyethylene glycol ethers of n-alcohols (lauryl, cetyl, stearyl and myristyl alcohol), liquid paraffin, lecithin oil, glycerol and the like. The applicants have combination of palmitate that the discovered stearate of ethyleneglycol and polyethylene glycol (Tefose 63), saturated polyglycolised glycerides M2130CS), glyceryl isostearate (Labrafil (isostearic peceol) and liquid paraffin proves very suitable for the implementation of the present invention. As a whole, the lipophilic excipients are present in a total proportion of from 10 to 40%, preferably from 30 to 35%, of the composition.
- 30 b) Mucoadhesive excipients: Cellulose polymers selected from sodium carboxymethylcellulose, hydroxypropyl methylcellulose, methylcellulose and the like, gelatin, colloidal anhydrous silica, or

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polyacrylic acid polymers, such as carbomers and polycarbophils. All these mucoadhesive excipients also possess gel-forming capacity. The applicants have discovered that the combination of polyacrylic cross-linked with divinyl glycol polycarbophil AA-1) and acrylic acid cross-linked with allyl esters of sucrose or pentaerythritol (e.g. carbopol 974P or carbopol 934P) proves very suitable for the implementation of the present invention. As a whole, the mucoadhesive excipients with gel-forming properties are present in a total proportion between 0.1 and 3%, preferably between 1 and 1.5% of the composition.

c) Preservatives: Parabens, such as methylparaben, butylparaben or propylparaben, benzoic acid, sorbic acid, boric acid and the like. As a whole, the preservatives are present in a total proportion between 0.01 and 0.3%, preferably between 0.1 and 0.2% of the composition.

Optionally, the compositions of the present invention can contain in addition suspending agents and humectants, such as povidone or propylene glycol, and neutralising agents for adjusting the viscosity of the composition, such as sodium hydroxide, triethanolamine (TEA) or ethylenediamine tetraacetic acid (EDTA). Povidone is normally used in concentrations of from 1 to 3% of the composition, preferably 2%. As for propylene glycol, it is normally used from 5 to 10% of the composition, preferably 7%.

Among the possible compositions, the invention relates preferably to creams and gels. For preparation of the

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compositions of the present invention, sertaconazole can be used as free base or in the form of a pharmaceutically acceptable salt. Among the pharmaceutically acceptable salts, the nitrate is preferred. The compositions may also contain mixtures of the free base with one or more salts as well as mixtures of two or more salts.

The cream formulations can be applied at two different concentrations of sertaconazole. The highest concentration applied inside the vagina and the is cream concentration is applied on the periphery of the infected The present invention relates more zone, the vulva. for internal application, cream precisely to the concentration The administered in a single dose. sertaconazole nitrate in this cream composition can range from 2 to 10%, and its quantity by volume can range from 4 to 6 ml. This corresponds to a dose of from 80 to 600 mg sertaconazole nitrate. According to one embodiment, the concentration is higher than 2 %, 3%, 4%, 5 % or 6% and, in particular, ranges from 3 to 10 %, or 4 to concentration of from 5 to 8%, and more preferably from 6 to 7% is preferred. A volume of 5 ml is preferred. This corresponds to a dose of from 250 to 400 mg and more preferably from 300 to 350 mg sertaconazole nitrate. For sertaconazole salts other than sertaconazole and nitrate the same concentrations apply. Alternatively, the present invention also relates to gels for application inside the vagina, which can be administered in a single dose. The concentration of the active ingredient in the gel formulations is similar to that of the corresponding cream formulations. However, the gels in contrast to the creams, do not necessarily contain lipophilic excipients in their proper administration of these formulation. For

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formulations (creams and gels), they can be conveniently packed inside an applicator, such as that described in ES 2,133,090, which constitutes one of the objects of the present invention. The size of the crystals of sertaconazole nitrate in the resulting cream should be below 80 μ m. Preferably, micronized sertaconazole nitrate having a particle size below 80 μ m is used.

On the other hand, the conventional cream for vulvar application mentioned in the preceding paragraph has a concentration of sertaconazole nitrate between 1 and 3%, preferably 2%. Its volume can range from 5 to 15 ml, preferably 10 ml. In the case of the cream or gel formulations for internal application, single or repeated doses can be administered. This cream composition is used for alleviating itching and irritation outside the vagina (in the vulva) in women infected with Candida spp. in both the vulva and the vagina, and represents a supplemental vaginal therapy with the concentrated cream or gel formulations, as described in the preceding paragraph.

Thus, a preferred embodiment of the present invention is a kit with the two formulations. The concentrated cream or gel formulation for internal use is conveniently packed in an applicator and prepared for its application in a single dose. The conventional cream for external use is packed in a conventional tube for its application in a single or repeated dose.

The release of sertaconazole nitrate from the two formulations, a concentrated intravaginal cream formulation (Example 1) and a conventional cream formulation, was tested. The formulation of the present invention releases

the active ingredient in a slow release profile, in contrast to the conventional cream formulation, which is also used for the treatment of the external area (vulva).

5 Unless indicated otherwise, concentrations and proportions given as percentage [%] refer to weight and thus mean "% by weight".

The present invention is further illustrated by - but not limited to - the following examples.

EXAMPLE 1: Preparation of 100 g of cream for intravaginal administration

15 Composition

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Micronised sertaconazole nitrate particle size below 80 µm	6.00 g
Tefose 63 ¹	20.00 g
Labrafil M 2130 CS ²	5.00 g
Isostearic peceol ³	2.00 g
Paraffin oil	8.00 g
Benzoic acid	0.10 g
Polycarbophil ⁴ AA-1	1.00 g
Carbopol 974 P ⁵	0.30 g
Purified water q.s. for	100.00 g

¹Tefose 63: Ethylene glycol and polyethylene glycol palmitate stearate

²Labrafil M 2130 CS: Saturated polyglycol glycerides

³Isoestearic peceol: Glyceryl isostearate

⁴Polycarbophil AA-1: Polyacrylic acid cross-linked with divinyl glycol ⁵Carbopol 974 P: Carbomer. Acrylic acid polymer cross-linked with sucrose and pentaerythritol allyl esters.

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Physicochemical properties

Appearance: White, odourless (or slight oily odour), semisolid cream of fluid consistency.

Penetrability: 43.5 ± 5% mm. 10

Viscosity: $347,000 \text{ cps} \pm 45\% (25^{\circ}\text{C})$.

EXAMPLE 2: In vitro dissolution and transdermal permeation tests

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The in vitro dissolution and transdermal permeation tests of sertaconazole nitrate from the cream formulations of Example 1 were evaluated in comparison with a conventional cream formulation of 2% sertaconazole nitrate.

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The composition for 100 g of the conventional cream is as follows:

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Micronised sertaconazole nitrate with particle size below 80 μm	2.00 g
Tefose 63 ¹	20.00 g
Labrafil MS 2230 ²	5.00 g
Isostearic peceol ³	2.00 g
Paraffin oil	8.00 g
Nipagin⁴	0.10 g
Sorbic acid	0.10 g

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Purified water q.s. for	100.00 g

¹Tefose 63: Ethylene glycol and polyethylene glycol palmitate stearate 2 Labrafil MS 2230: Saturated ($C_{10}-C_{18}$) polyoxyethylene glycol and glycol glycerides

³Isoestearic peceol: Glyceryl isostearate

⁴Nipagin: Methyl p-hydroxybenzoate.

Both tests were carried out with Franz cell-type diffusion systems, with a diffusional area of $2.54~\rm cm^2$. 1 ml of cream was placed in the donor compartment and 11 ml of a suitable receptor medium were placed in the receptor compartment. For the dissolution test, a $0.45-\mu m$ Millipore membrane of nylon esters was used, and the receptor medium was made up of a mixture of ethanol-water (1:1). Vaginal epithelium was used as permeation membrane and phosphate buffer solution at pH 7.4 was used as the receptor medium.

The 4-cm² membranes used in the permeation test were formed by reconstituted vaginal epithelial cells (5-day culture) from transformed cells of human vaginal epithelium on polycarbonate support. These cells were obtained from cell lines of vulvar epidermoid carcinomas. The test temperature was 32°C for both cases.

In the light of the physicochemical properties of sertaconazole, it can be assumed that the maximum quantity permeated is about 1% of the quantity located on the membrane. Under such assumption, the maximum quantity of sertaconazole that reaches the receptor compartment is 6.18 g/ml.

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The curves showing the release of sertaconazole nitrate from the two cream formulations are plotted in Fig. 1. Starting from 5 ml of the cream formulation of Example 1, 125 mg of the active ingredient have already been released at 24 hours and on the following days at the rate of 50-60 mg/day. Thus, it is observed that the delivery of the active ingredient is 81% after 5 days.

Moreover, the *in vitro* permeation test shows that the active substance present in the vaginal cream formulation of Example 1 permeates less than 0.1% of the dose.

The toxicity study performed to assess the non-clinical vaginal tolerance of the mucoadhesive cream proves a good tolerance after being applied at single or repeated doses to rats (method according to CMP/SWP/21H5/00).

EXAMPLE 3: Preparation of 100 g of gel for intravaginal administration

Starting from the appropriate components and according to standard procedures of pharmaceutical technology, the following gel composition was obtained:

Micronised sertaconazole nitrate with particle size below 80 μm	6.00 g
Carbopol 974 P ¹	0.70 g
Polycarbophil AA-1 ²	0.30 g
Propylene glycol	7.00 g
Nipagin ³	0.16 g
Nipasol⁴	0.04 g

Povidone	2.00 g
TEA ⁵	*
Purified water q.s. for	100.00 g

¹Carbopol 974 P: Carbomer. Acrylic acid polymer cross-linked with sucrose and pentaerythritol allyl esters.

²Polycarbophil AA-1: Polyacrylic acid cross-linked with divinyl glycol

³Nipagin: Methyl *p*-hydroxybenzoate ⁴Nipasol: Propyl *p*-hydroxybenzoate

⁵TEA: Triethanolamine

* Quantity sufficient to adjust the viscosity